

## STIC Search Report Biotech-Chem Library

## STIC Database Tracking Number 2002

TO: Shirley Gembeh

Location: rem/3A44/3C70

Art Unit: 1614

**Tuesday, May 30, 2006** 

Case Serial Number: 10/743354

From: Deirdre Arnold

**Location: Biotech-Chem Library** 

**REM 1A55** 

Phone: 571-272-2532

Deirdre.Arnold@uspto.gov

## Search Notes



- Please check the structure for accuracy.
- In accordance with your request, only the elected species (circled) was searched. There were very few hits. If you would like to broaden the search by making the structure less defined, please contact me.
- Beware of false hits on the names in the inventor search.

Please feel free to contact me if you have any questions or would like to amend the search.

Thank you for using STIC services.

Regards,

Deirdre Arnold



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Scientific and Technical Information Center

Searcher Prep & Review Time: \_\_\_\_

Online Time:

May 29 22.6 SI	EARCH REC	QUEST FORM
	umber: 2-3504	Examiner #: 80889 Date: 5/30/06  Serial Number: 10,743,354
Location (Bldg/Room#): Rem 3144 (Ma	ailbox #): <u>3C 70</u>	Results Format Preferred (circle): PAPER DISK
To ensure an efficient and quality search, ples		over sheet, claims, and abstract or fill out the following:
Title of Invention: Helcoar	ylallanoic C	acids as integrin receptor.
Inventors (please provide full names):	Boys et a	٠
Earliest Priority Date: 12/20/0	12	
elected species or structures, keywords, synonyn	ms, acronyms, and registr	pecifically as possible the subject matter to be searched. Include the y numbers, and combine with the concept or utility of the invention. levant citations, authors, etc., if known.
*For Sequence Searches Only* Please include	all pertinent information	(parent, child, divisional, or issued patent numbers) along with the
appropriate serial number.	alasked	(parent, child, divisional, or issued patent numbers) along with the  empound (circled Cmp)  pplicants response)  y possible)
Docse Search	~ election	relicants reaponse)
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See our	/A-8AP	y posson)
Thank you		
·		
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## ### ##############################		
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******	*****	***********
STAFF USE ONLY	Type of Search	Vendors and cost where applicable
Searcher:	NA Sequence (#	
Searcher Phone #:	AA Sequence (#	Westlaw WWW/Internet
Searcher Location:	Bibliographic	In-house sequence systems
Date Searcher Picked Up:	Биловіаріне	
Date Completed:	Litigation	Interference SPDI Encode/Transl

Fulltext Other



# STIC SEARCH RESULTS FEEDBACK FORM

## Biotech:Chem Library

Questions about the scope or the results of the search? Contact the searcher or contact:

Mary Hale, Information Branch Supervisor 571-272-2507 Remsen E01 D86

O	Untary Results Feedback Form
>	I am an examiner in Workgroup: Example: 1610
۶	Relevant prior art found, search results used as follows:
	☐ 102 rejection
	☐ 103 rejection
	Cited as being of interest.
	Helped examiner better understand the invention.
	Helped examiner better understand the state of the art in their technology.
	Types of relevant prior art found:
	☐ Foreign Patent(s)
	<ul> <li>Non-Patent Literature         (journal articles, conference proceedings, new product announcements etc.)</li> </ul>
>	Relevant prior art not found:
	Results verified the lack of relevant prior art (helped determine patentability).
	Results were not useful in determining patentability or understanding the invention.
Co	omments:

Drop off or send completed forms to Stile/Biotech-Chem Library Remsen Bldg



#### What is claimed is:

- 1. A pharmaceutical composition comprising a pharmaceutically acceptable carrier and a compound selected from the group consisting of: 3-(3,5-ditert-butylphenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid (TFA salt);
- 5 3-(3-tert-butyl-5-iodophenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
  - 3-(3-tert-butyl-5-bromophenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
  - 3-(5-tert-Butyl-2-hydroxyphenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-
- 10 yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
  - 3-[3,5-Ditert-butyl-2-(carboxymethoxy)phenyl]-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
  - 3-(5-tert-Butyl-2-methoxyphenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
- 3-(3,5-Ditert-butyl-4-methoxyphenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
  - 3-{3-tert-Butyl-5-[2,2,2-trifluoro-1-hydroxy-1-(trifluoromethyl)ethyl]-phenyl}-4-{3-[3-
  - (5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid;
  - $3-(3,4-Dichlorophenyl)-4-\{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-1,2,4-1,2,3-1,2,4-1$
- 20 oxadiazol-5-yl}butanoic acid trifluoroacetate;
  - 3-(3-Fluoro-4-methylphenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid hydrochloride;
  - 3-(4-Phenoxyphenyl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid trifluoroacetate;
- 3-(1-Benzofuran-2-yl)-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}butanoic acid trifluoroacetate;
  - 3-[4-(Benzyloxy)phenyl]-4-{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-
  - 1,2,4-oxadiazol-5-yl}butanoic acid trifluoroacetate;
  - $3-[4-(Methylsulfonyl)phenyl]-4-\{3-[3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyl]-4-\{3-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)propyllapro-1,8-(5,6,7,8-tetrahydro-1,8-(5,6,7,8-tetrahydro-1,8-(5,6,7,8-tetrahydro-1,8-(5,6,7,8-tetrahydro-1,8-(5,6,7,8-tetrahydro-1,8-(5,6,7,8-tetra$
- 30 1,2,4-oxadiazol-5-yl}butanoic acid trifluoroacetate;
  - 4-{3-[3-(5,6,7,8-Tetrahydro-1,8-naphthyridin-2-yl)propyl]-1,2,4-oxadiazol-5-yl}-3-[4-(trifluoromethoxy)phenyl]butanoic acid trifluoroacetate;



### UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Vignia 22313-1450 www.uspto.gov

## 

Bib Data Sheet

**CONFIRMATION NO. 9317** 

Dib Data Officet											
<b>SERIAL NUMBER</b> 10/743,354											
Michael B. Tolle Nizal Samuel Cl Ish Kumar Khan Maria Nguyen, A Victoria L. Dowr Scott B. Mohler, Glen J. Gesicki, Thomas D. Pen Barbara B. Cher Yaping Wang, A Albert Khilevich, Bipinchandra N. Yi Yu, Glenview John A. Wendt, Heather Stenma Hongwei Wu, Br Renee M. Huff, Srinivasan Raj N Balekudru Deva Hwang-Fun Lu, Mark Russell, G Dale P. Spangle Mihir D. Parikh,  ** CONTINUING DAT This appln claim ** FOREIGN APPLICA	nan, East Hanover, NJ; fson, Dardenne Prairie handrakumar, Grafton, ina, Libertyville, IL; Ann Arbor, MI; is, Pinckney, MI; Chicago, IL; Chicago, IL; ning, Elmhurst, IL; n, Glenview, IL; acton, MA; , Buffalo Grove, IL; Desai, Vernon Hills, IL; , IL; South Lyon, MI; ark, Chicago, IL; uffalo Grove, IL; Park Ridge, IL; Nagarajan, Chesterfield, MO; Bullwin, MO; Burnee, IL; er, San Diego, CA; Chesterfield, MO;  A ***********************************	, MO; MA; d, MO; 12/20/2002									
Foreign Priority claimed 35 USC 119 (a-d) conditions met Verified and Acknowledged Exam		fter STATE OF COUNTRY		TOTAL CLAIMS 5	INDEPENDENT CLAIMS 1						
<b>ADDRESS</b> 28940											
TITLE											
Heteroarylalkanoic aci	ds as integrin receptor	antagonists derivative	es								

=> d que stat 19

L1 1 SEA FILE=HCAPLUS ABB=ON PLU=ON US2003-743354/APPS

L3 TRANSFER PLU=ON L1 1- RN : 387 TERMS

L4 387 SEA FILE=REGISTRY ABB=ON PLU=ON L3

L5 131 SEA FILE=REGISTRY ABB=ON PLU=ON L4 AND ?NAPHTHYRIDIN?/CNS

L9 1 SEA FILE=REGISTRY ABB=ON PLU=ON L5 AND I/ELS

=> d ide 19

YOU HAVE REQUESTED DATA FROM FILE 'REGISTRY' - CONTINUE? (Y) /N:y

L9 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2006 ACS on STN

RN 724770-27-8 REGISTRY

ED Entered STN: 10 Aug 2004

CN 1,2,4-Oxadiazole-5-butanoic acid, β-[3-(1,1-dimethylethyl)-5-iodophenyl]-3-[3-(1,5,6,7-tetrahydro-1,8-naphthyridin-2-yl)propyl]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C27 H33 I N4 O3

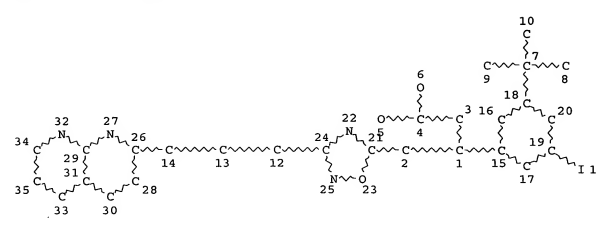
SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

- 1 REFERENCES IN FILE CA (1907 TO DATE)
- 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> => d que stat 116 L14 . STR



Page 1-A

11

Page 1-B

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 35

STEREO ATTRIBUTES: NONE

L16 1 SEA FILE=REGISTRY SS FUL L14

100.0% PROCESSED SEARCH TIME: 00.00.01 3 ITERATIONS

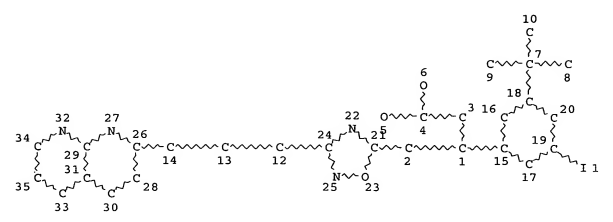
1 ANSWERS

=> d que	nos 117	
L1	1 :	SEA FILE=HCAPLUS ABB=ON PLU=ON US2003-743354/APPS
L3	•	TRANSFER PLU=ON L1 1- RN : 387 TERMS
L4	387	SEA FILE=REGISTRY ABB=ON PLU=ON L3
L5	131 :	SEA FILE=REGISTRY ABB=ON PLU=ON L4 AND ?NAPHTHYRIDIN?/CNS
Ь9	1 :	SEA FILE=REGISTRY ABB=ON PLU=ON L5 AND I/ELS
L14	:	STR
L16	1 :	SEA FILE=REGISTRY SSS FUL L14
L17	0 :	SEA FILE=REGISTRY ABB=ON PLU=ON L16 NOT L9

=> d que nos 124

L1	1	SEA FILE=HCAPLUS ABB=ON PLU=ON US2003-743354/APPS
L3		TRANSFER PLU=ON L1 1- RN : 387 TERMS
L4	387	SEA FILE=REGISTRY ABB=ON PLU=ON L3
L5	131	SEA FILE=REGISTRY ABB=ON PLU=ON L4 AND ?NAPHTHYRIDIN?/CNS
L9	1	SEA FILE=REGISTRY ABB=ON PLU=ON L5 AND I/ELS
L23	1	SEA FILE=REGISTRY ABB=ON PLU=ON 724770-27-8/RN,CRN
L24	0	SEA FILE=REGISTRY ABB=ON PLU=ON L23 NOT L9

=> d que stat 118 L14



Page 1-A

11

L14

Page 1-B NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 35

STEREO ATTRIBUTES: NONE

L18 0 SEA FILE=BEILSTEIN SSS(FUL) L14

100.0% PROCESSED 0 ITERATIONS

STR

0 ANSWERS SEARCH TIME: 00.00.02

=> d que stat 120

10 32 29 14 12 13 2 17 33 30

Page 1-A

11

Page 1-B NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

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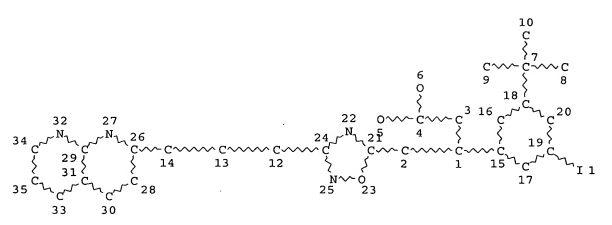
STEREO ATTRIBUTES: NONE

L20 0 SEA FILE=CHEMINFORMRX SSE FULL L14 ( 0 REACTIONS)

100.0% DONE 0 VERIFIED 0 HIT RXNS 0 DOCS

SEARCH TIME: 00.00.02

=> d que stat 122 L14 STR



Page 1-A

11

Page 1-B NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 35

STEREO ATTRIBUTES: NONE

L22 0 SEA FILE=MARPAT SSS(FUL)L14

100.0% PROCESSED 42 ITERATIONS

0 ANSWERS

SEARCH TIME: 00.00.01

#### => d his 120-132

L23

FILE 'STNGUIDE' ENTERED AT 12:20:12 ON 30 MAY 2006

FILE 'MARPAT' ENTERED AT 12:20:15 ON 30 MAY 2006

L21 0 S L14 SAM L22 0 S L14 FUL

SAVE TEMP L22 GEM354MAR1/A

FILE 'STNGUIDE' ENTERED AT 12:21:44 ON 30 MAY 2006

FILE 'REGISTRY' ENTERED AT 12:23:18 ON 30 MAY 2006

1 S 724770-27-8/RN,CRN

L24 0 S L23 NOT L9

FILE 'STNGUIDE' ENTERED AT 12:23:45 ON 30 MAY 2006

FILE 'WPIX' ENTERED AT 12:24:13 ON 30 MAY 2006 SELECT L2 1- DCRE

L25 98 S E13-E110/DCSE

FILE 'STNGUIDE' ENTERED AT 12:25:32 ON 30 MAY 2006

FILE 'WPIX' ENTERED AT 12:26:09 ON 30 MAY 2006

L26 1 S L25 AND (C27 H33 I N4 O3)/MF

SELECT L26 1- DCSE

L27 1 S E111/KW

L28 0 S L14 SAM L29 1 S L14 FUL

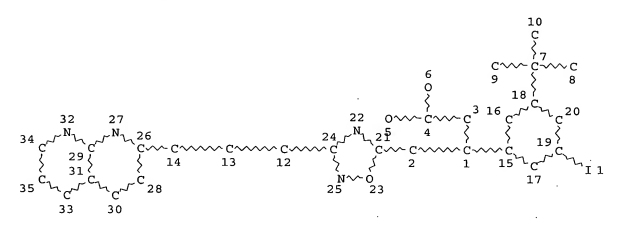
L29 1 S L14 FUL L30 1 S L29/DCR

SELECT L29 1- SDCN

L31 1 S E112/DCN

L32 1 S L27 OR L30 OR L31

=> d que stat 132 L14 STR



Page 1-A

11

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Page 1-B
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
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GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 35

#### => d his 111

(FILE 'HCAPLUS, TOXCENTER, USPATFULL' ENTERED AT 12:11:11 ON 30 MAY 2006) L11 3 S L9

=> d que stat l11

L1 1 SEA FILE=HCAPLUS ABB=ON PLU=ON US2003-743354/APPS
L3 TRANSFER PLU=ON L1 1- RN : 387 TERMS
L4 387 SEA FILE=REGISTRY ABB=ON PLU=ON L3
L5 131 SEA FILE=REGISTRY ABB=ON PLU=ON L4 AND ?NAPHTHYRIDIN?/CNS
L9 1 SEA FILE=REGISTRY ABB=ON PLU=ON L5 AND I/ELS
L11 3 SEA L9

=> dup rem 111 118 120 122 132 L18 HAS NO ANSWERS

L20 HAS NO ANSWERS

L22 HAS NO ANSWERS

DUPLICATE IS NOT AVAILABLE IN 'BEILSTEIN, CHEMINFORMRX'.

ANSWERS FROM THESE FILES WILL BE CONSIDERED UNIQUE
FILE 'HCAPLUS' ENTERED AT 13:00:18 ON 30 MAY 2006
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

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FILE 'TOXCENTER' ENTERED AT 13:00:18 ON 30 MAY 2006 COPYRIGHT (C) 2006 ACS

FILE 'USPATFULL' ENTERED AT 13:00:18 ON 30 MAY 2006 CA INDEXING COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'WPIX' ENTERED AT 13:00:18 ON 30 MAY 2006
COPYRIGHT (C) 2006 THE THOMSON CORPORATION
PROCESSING COMPLETED FOR L11
PROCESSING COMPLETED FOR L20
PROCESSING COMPLETED FOR L22
PROCESSING COMPLETED FOR L32
L68
2 DUP REM L11 L18 L20 L22 L32 (2 DUPLICATES REMOVED)
ANSWER '1' FROM FILE HCAPLUS

#### ANSWER '2' FROM FILE USPATFULL

=> file stnguide FILE 'STNGUIDE' ENTERED AT 13:00:24 ON 30 MAY 2006 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY, JAPAN SCIENCE AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: May 26, 2006 (20060526/UP).

=> d ibib ed ab ind hitstr
YOU HAVE REQUESTED DATA FROM FILE 'HCAPLUS, USPATFULL' - CONTINUE? (Y)/N:y

L68 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1 ACCESSION NUMBER: 2004:565087 HCAPLUS DOCUMENT NUMBER: 141:123408 Heteroarylalkanoic acids as integrin receptor TITLE: antagonists Boys, Mark L.; Schretzman, Lori A.; Tollefson, Michael INVENTOR (S): B.; Chandrakumar, Nizal S.; Khanna, Ish K.; Nguyen, Maria; Downs, Victoria; Mohler, Scott B.; Gesicki, Glen J.; Penning, Thomas D.; Chen, Barbara B.; Wang, Yaping; Khilevich, Albert; Desai, Bipinchandra N.; Yu, Yi; Wendt, John A.; Stenmark, Heather; Wu, Lisa; Huff, Renee M.; Nagarajan, Srinivasan R.; Devadas, Balekudru; Lu, Hwang-fun; Russell, Mark; Spangler, Dale P.; Parikh, Mihir D.; Clare, Michael PATENT ASSIGNEE(S): Pharmacia Corporation, USA SOURCE: PCT Int. Appl., 266 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -----------WO 2004058254 A1 20040715 WO 2003-US40898 20031222 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW

RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GO, GW, MI, MP, NE, SN, TD, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, CA 2507699 AA 20040715 CA 2003-2507699 20031222 AU 2003299807 **A1** 20040722 AU 2003-299807 20031222 US 2005043344 **A**1 20050224 US 2003-743354 20031222 EP 1592421 A1 20051109 EP 2003-800081 20031222 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK BR 2003017600 Α 20051129 BR 2003-17600 20031222 PRIORITY APPLN. INFO.: US 2002-435467P P 20021220 WO 2003-US40898 W 20031222

OTHER SOURCE(S): MARPAT 141:123408

ED Entered STN: 15 Jul 2004

AB The present invention relates to pharmaceutical compns. comprising compds. I [A = (un)saturated and/or (un)substituted 4-8 membered monocyclic or 7-12 membered bicyclic ring, containing 1 to 5 heteroatoms selected from the group consisting of O, N or S; ring A may further contain a carboxamide, sulfone, sulfonamide, or an acyl group; A1 = (un)saturated and/or (un)substituted 5-9 membered monocyclic or 8-14 membered polycyclic heterocycle containing at least one N; or A1 = substituted urea, iminourea or thiourea alicyclic or cyclic analog; Z1 = CH2, CH2O, O, NH, CO, S, SO, CHOH, SO2; Z2 = (un)substituted 1-5 carbon linker optionally containing one or

Gembeh 10/743,354 05/30/2006 more heteroatoms; alternatively Z1-Z2 may contain a carboxamide, sulfone, sulfonamide, alkenyl, acyl group, or aryl or heteroaryl ring; X = CO, SO2, S, O, substituted amine, substituted CH; Y = CO, SO2, substituted amine, etc.; Y5 = C or N; Y3 and Y4 independently = H, halo, (un)substitutedalkyl, -aryl, -alkene, etc.; or Y3 and Y4 together form a (un)saturated and/or (un) substituted 3-8 membered monocyclic or a 7-11 membered bicyclic ring optionally containing heteroatoms; or X and Y3 form a 3-7 membered monocyclic ring optionally containing heteroatoms; Rb = OH, alkoxy, arylamine, etc.], or a pharmaceutically acceptable salt thereof, methods of selectively inhibiting or antagonizing the  $\alpha\nu\beta3$  and/or the ανβ5 integrin without significantly inhibiting the  $\alpha\nu\beta6$  integrin, and methods to prepare I. Thus, e.g., II was prepared in four steps with oxadiazole ring forming via cyclization reaction of amide oxime III with cyclic anhydride IV (preparation given). I antagonize  $\alpha v\beta 3$  integrin with an IC50 values ranging from 0.1 nM to 100  $\mu M$  in the 293-cell assay. Similarly, I also antagonized  $\alpha\nu\beta5$  integrin with an IC50 values of < 50  $\mu M$  in the cell adhesion assay. ICM A61K031-4245 ICS A61P019-02; A61P019-10; A61P027-00; A61P035-00; A61P035-04; A61P043-00 23-16 (Aliphatic Compounds) Section cross-reference(s): 1, 28, 63 butanoic acid heteroaryl deriv prepn integrin receptor antagonist; alkanoic acid heteroaryl deriv prepn integrin receptor antagonist Neoplasm (humoral hypercalcemia of malignancy; preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha \nu \beta 3$ and/or ανβ5 integrin receptor) Eye, disease (macula, degeneration; preparation of heteroaryl butanoic acid derivs. as

IT

selective inhibitors or antagonists of  $\alpha v\beta 3$  and/or ανβ5 integrin receptor)

IT Neoplasm

IC

CC

ST

ΙT

(metastasis; preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha\nu\beta3$  and/or  $\alpha\nu\beta5$ integrin receptor)

Angiogenesis IT

Antiarteriosclerotics

Antiarthritics

Antitumor agents

Arthritis

Atherosclerosis

Drug delivery systems

Osteoporosis

(preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha v\beta 3$  and/or  $\alpha v\beta 5$  integrin receptor)

IT Artery, disease

> (restenosis; preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha v\beta 3$  and/or  $\alpha v\beta 5$ integrin receptor)

IT Eye, disease

(retinopathy; preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha v \beta 3$  and/or  $\alpha v \beta 5$ integrin receptor)

IT Cell migration

> (smooth muscle; preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha v\beta 3$  and/or  $\alpha v\beta 5$ integrin receptor)

```
IT
     Integrins
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (ανβ3; preparation of heteroaryl butanoic acid derivs. as
        selective inhibitors or antagonists of \alpha v\beta 3 and/or
        ανβ5 integrin receptor)
IT
     Integrins
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (ανβ5; preparation of heteroaryl butanoic acid derivs. as
        selective inhibitors or antagonists of αvβ3 and/or
        ανβ5 integrin receptor)
     724769-51-1P
IT
     RL: BPN (Biosynthetic preparation); PAC (Pharmacological activity); SPN
     (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study);
     PREP (Preparation); USES (Uses)
        (preparation of heteroaryl butanoic acid derivs. as selective inhibitors or
        antagonists of \alpha v\beta 3 and/or \alpha v\beta 5 integrin
        receptor)
ΙT
     17004-92-1P
     RL: BPN (Biosynthetic preparation); PUR (Purification or recovery); RCT
     (Reactant); BIOL (Biological study); PREP (Preparation); RACT (Reactant or
        (preparation of heteroaryl butanoic acid derivs. as selective inhibitors or
        antagonists of \alpha v\beta 3 and/or \alpha v\beta 5 integrin
        receptor)
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     or reagent)
        (preparation of heteroaryl butanoic acid derivs. as selective inhibitors or
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        receptor)
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        receptor)
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     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (preparation of heteroaryl butanoic acid derivs. as selective inhibitors or
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       receptor)
IT
     67-36-7, 4-Phenoxybenzaldehyde
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     Thiosemicarbazide
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                    105-58-8, Diethylcarbonate
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                    110-89-4, Piperidine, reactions
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                                                       110-91-8, Morpholine,
     reactions
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                    351-54-2, 3-Fluoro-4-methoxybenzaldehyde
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                          656-42-8, 2,2-Difluoro-1,3-benzodioxole-5-
                      659-28-9, 4-Trifluoromethoxybenzaldehyde
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     867-13-0, Triethylphosphonoacetate
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    Benzenecarbothioamide
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    hydriodide
                  5693-62-9
                              5717-37-3, (Carbethoxyethylidene) triphenylphospho
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     4-Methyl-3-thiosemicarbazide
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     7311-34-4, 3,5-Dimethoxybenzaldehyde
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    N-(2-hydroxyethyl)carbamate 29335-36-2
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    RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of heteroaryl butanoic acid derivs. as selective inhibitors or
       antagonists of \alpha v\beta 3 and/or \alpha v\beta 5 integrin
       receptor)
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha\nu\beta3$  and/or  $\alpha\nu\beta5$  integrin receptor)

#### IT 724770-27-8P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha\nu\beta3$  and/or  $\alpha\nu\beta5$  integrin receptor)

#### RN 724770-27-8 HCAPLUS

CN 1,2,4-Oxadiazole-5-butanoic acid, β-[3-(1,1-dimethylethyl)-5iodophenyl]-3-[3-(1,5,6,7-tetrahydro-1,8-naphthyridin-2-yl)propyl]- (9CI)
(CA INDEX NAME)

=> d ibib ab hitstr 2
YOU HAVE REQUESTED DATA FROM FILE 'HCAPLUS, USPATFULL' - CONTINUE? (Y)/N:y

L68 ANSWER 2 OF 2 USPATFULL on STN

ACCESSION NUMBER: 2005:50544 USPATFULL

Heteroarylalkanoic acids as integrin receptor TITLE:

antagonists derivatives

INVENTOR (S): Boys, Mark L., Brighton, MI, UNITED STATES

> Schretzman, Lori A., East Hanover, NJ, UNITED STATES Tollefson, Michael B., Dardenne Prairie, MO, UNITED

STATES

Chandrakumar, Nizal Samuel, Grafton, MA, UNITED STATES Khanna, Ish Kumar, Libertyville, IL, UNITED STATES

Nguyen, Maria, Ann Arbor, MI, UNITED STATES Downs, Victoria L., Pinckney, MI, UNITED STATES Mohler, Scott B., Chicago, IL, UNITED STATES Gesicki, Glen J., Chicago, IL, UNITED STATES Penning, Thomas D., Elmhurst, IL, UNITED STATES

Chen, Barbara B., Glenview, IL, UNITED STATES Wang, Yaping, Acton, MA, UNITED STATES

Khilevich, Albert, Buffalo Grove, IL, UNITED STATES Desai, Bipinchandra N., Vernon Hills, IL, UNITED STATES

Yu, Yi, Glenview, IL, UNITED STATES

Wendt, John A., South Lyon, MI, UNITED STATES Stenmark, Heather, Chicago, IL, UNITED STATES Wu, Hongwei, Buffalo Grove, IL, UNITED STATES Huff, Renee M., Park Ridge, IL, UNITED STATES

Nagarajan, Srinivasan Raj, Chesterfield, MO, UNITED

STATES

Devadas, Balekudru, Chesterfield, MO, UNITED STATES

Lu, Hwang-Fun, Ballwin, MO, UNITED STATES Russell, Mark, Gurnee, IL, UNITED STATES

Spangler, Dale P., San Diego, CA, UNITED STATES Parikh, Mihir D., Chesterfield, MO. UNITED STATES

PATENT ASSIGNEE(S): Pharmacia Corporation (U.S. corporation)

> NUMBER KIND DATE -----

PATENT INFORMATION:

US 2005043344 A1 US 2003-743354 A1 20050224

APPLICATION INFO.: 20031222 (10)

> NUMBER DATE -----

PRIORITY INFORMATION:

US 2002-435467P 20021220 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

SENNIGER POWERS LEAVITT AND ROEDEL, ONE METROPOLITAN

SQUARE, 16TH FLOOR, ST LOUIS, MO, 63102

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: LINE COUNT: 6480

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to pharmaceutical compositions comprising compounds of the Formula I, or a pharmaceutically acceptable salt

thereof, and methods of selectively inhibiting or antagonizing the  $\alpha.sub.V\beta.sub.3$  and/or the  $\alpha.sub.V\beta.sub.5$  integrin

without significantly inhibiting the  $\alpha.sub.V\beta.sub.6$  integrin.

IT 724770-27-8P

(preparation of heteroaryl butanoic acid derivs. as selective inhibitors or antagonists of  $\alpha v\beta 3$  and/or  $\alpha v\beta 5$  integrin receptor)

RN 724770-27-8 USPATFULL

CN 1,2,4-Oxadiazole-5-butanoic acid, β-{3-(1,1-dimethylethyl)-5iodophenyl]-3-[3-(1,5,6,7-tetrahydro-1,8-naphthyridin-2-yl)propyl}(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & t-Bu \\ & & & & \\ & & & \\ N & & & \\ & & & \\ N & & \\ \end{array}$$

#### => d his 167

(FILE 'HCAPLUS, MEDLINE, BIOSIS, PASCAL, JICST-EPLUS, CABA, LIFESCI, DRUGU, DRUGB, VETU, VETB, WPIX, SCISEARCH, CONF, CONFSCI, DISSABS' ENTERED AT 12:37:36 ON 30 MAY 2006)

L67 11 DUP REM L66 (10 DUPLICATES REMOVED)

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L64
L65
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L66
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             11 DUP REM L66 (10 DUPLICATES REMOVED)
1.67
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=> d ibib ed ab 167 1-11
YOU HAVE REQUESTED DATA FROM FILE 'HCAPLUS, BIOSIS' - CONTINUE? (Y)/N:y

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L67 ANSWER 1 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1
ACCESSION NUMBER: 2006:16564 HCAPLUS
DOCUMENT NUMBER: 144:254037
TITLE: Synthesis of 2,5-thiazole butanoic acids as potent and selective ανβ3 integrin receptor antagonists with improved oral pharmacokinetic properties
AUTHOR(S): Wendt, John A.; Wu, Hongwei; Stenmark,
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Heather G.; Boys, Mark L.; Downs,
                          Victoria L.; Penning, Thomas D.; Chen,
                          Barbara B.; Wang, Yaping; Duffin, Tiffany; Finn, Mary
                          Beth; Keene, Jeffery L.; Engleman, V. Wayne; Freeman,
                          Sandra K.; Hanneke, Melanie L.; Shannon, Kristen E.;
                          Nickols, Maureen A.; Steininger, Christina N.;
                          Westlin, Marissa; Klover, Jon A.; Westlin, William;
                          Nickols, G. Allen; Russell, Mark A.
CORPORATE SOURCE:
                         Department of Medicinal Chemistry, Pfizer Global
                          Research and Development, Ann Arbor, MI, 48105, USA
                          Bioorg. Med. Chem. Lett. (2006), 16(4), 845-849
SOURCE:
                          CODEN: BMCLE8; ISSN: 0960-894X
PUBLISHER:
                          Elsevier B.V.
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                         English
OTHER SOURCE(S):
                         CASREACT 144:254037
     Entered STN: 08 Jan 2006
     A series of 2,5-thiazoles, e.g. I [R = 3-FC6H4, 3,4-(MeO) 2C6H3,
AB
     6-methoxy-3-pyridyl, 2-phenyl-5-thiazolyl, etc.], which are potent
     antagonists of the integrin \alpha v\beta 3 and show
     selectivity relative to the other integrins, such as
     \alpha IIb\beta 3 and \alpha \nu \beta 6, has been synthesized. These
     analogs were demonstrated to have high bioavailability relative to other
     relative heterocyclic analogs.
                                THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                         26
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L67 ANSWER 2 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 2
ACCESSION NUMBER:
                         2006:16560 HCAPLUS
DOCUMENT NUMBER:
                         144:254069
TITLE:
                         Convergent, parallel synthesis of a series of
                         β-substituted 1,2,4-oxadiazole butanoic
                         acids as potent and selective \alpha v\beta 3 receptor
                         antagonists
AUTHOR (S):
                         Boys, Mark L.; Schretzman, Lori A.
                         ; Chandrakumar, Nizal S.; Tollefson,
                         Michael B.; Mohler, Scott B.;
                         Downs, Victoria L.; Penning, Thomas D.
                          ; Russell, Mark A.; Wendt, John A.; Chen,
                         Barbara B.; Stenmark, Heather G.; Wu,
                         Hongwei; Spangler, Dale P.; Clare, Michael;
                         Desai, Bipin N.; Khanna, Ish K.;
                         Nguyen, Maria N.; Duffin, Tiffany; Engleman, V. Wayne;
                         Finn, Mary Beth; Freeman, Sandra K.; Hanneke, Melanie
                         L.; Keene, Jeffery L.; Klover, Jon A.; Nickols, G.
                         Allen; Nickols, Maureen A.; Steininger, Christina N.;
                         Westlin, Marisa; Westlin, William; Yu, Yi X.; Wang,
                         Yaping; Dalton, Christopher R.; Norring, Sarah A.
CORPORATE SOURCE:
                         Department of Chemistry, PfizerGlobal Research and
                         Development, Ann Arbor, MI, 48105, USA
                         Bioorg. Med. Chem. Lett. (2006), 16(4), 839-844
SOURCE:
                         CODEN: BMCLE8; ISSN: 0960-894X
                         Elsevier B.V.
PUBLISHER:
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
OTHER SOURCE(S):
                         CASREACT 144:254069
    Entered STN: 08 Jan 2006
     A series of 1,2,4-oxadiazoles, e.g. I [R1 = H, R2 = H, Me, HC.tplbond.C,
     Ph, 3-pyridyl, 2-methyl-5-thiazolyl, etc.; R1 = Me, R2 = Me, Ph,
     3-pyridyl; R1R2 = (CH2)4], which are potent antagonists of the
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integrin  $\alpha\nu\beta3$  and, in addition, show selectivity relative to the other  $\beta3$  integrin  $\alpha IIb\beta3$ , has been synthesized. In whole cells, the majority of these analogs also demonstrated modest selectivity against other  $\alpha\nu$  integrins such as  $\alpha\nu\beta1$  and  $\alpha\nu\beta6$ .

REFERENCE COUNT:

THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L67 ANSWER 3 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 3

ACCESSION NUMBER: 2004:565087 HCAPLUS

DOCUMENT NUMBER: 141:123408

TITLE: Heteroarylalkanoic acids as integrin

receptor antagonists.

INVENTOR(S): Boys, Mark L.; Schretzman, Lori A.

; Tollefson, Michael B.; Chandrakumar,

Nizal S.; Khanna, Ish K.; Nguyen, Maria; Downs, Victoria; Mohler,

Scott B.; Gesicki, Glen J.;

Penning, Thomas D.; Chen, Barbara B.; Wang, Yaping; Khilevich, Albert; Desai, Bipinchandra N.; Yu, Yi; Wendt, John A.; Stenmark, Heather; Wu, Lisa; Huff, Renee M.; Nagarajan, Srinivasan R.; Devadas, Balekudru; Lu, Hwang-fun; Russell, Mark; Spangler,

Dale P.; Parikh, Mihir D.; Clare,

Michael

PATENT ASSIGNEE(S):

SOURCE:

Pharmacia Corporation, USA PCT Int. Appl., 266 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

1	PATENT NO.			KIND DATE			APPLICATION NO.					DATE							
v	WO 2004058254			A1 20040715			WO 2003-US40898						20	0031	222				
	1	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
			•	•	•			DE,				-							
			•	•	•	•		ID,		•		•							
			•	•	•	•		LV,		•	٠.	•				•	•		
			•	•	•			PT,	•	•		-	-	•		•	•		
			•	•		•	•	UA,	•	•	•	•	•		-	•		/	
		pw.		•		-		MW,		-	-							Δ7.	
		,				-	-	ΤJ,		-		•					-	-	
				•		-		HU,			•	-							
			•	•	•			CI,		•	•	•	•						TC
,	ר מי	E07				•		•		•	•						•	•	10
									CA 2003-2507699 AU 2003-299807										
	-																		
Ţ	US 2	005	0433	44		A1		2005	0224		US 2	003-	7433	54		20	0031	222	
1	EP 1	5924	421			A1		2005	1109	:	EP 2	003-	80008	31		20	0031	222	
		R:	AT,	BE,	CH,	DÉ,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK		
I	• • • • • • • • • • • • • • • • • • • •				Α		2005	1129	BR 2003-17600										
PRIOR	PRIORITY APPLN. INFO.:			. :									57P	P 20021220					
										WO 2003-US40898					W 20031222				
											-								

OTHER SOURCE(S): MARPAT 141:123408

ED Entered STN: 15 Jul 2004

AB The present invention relates to pharmaceutical compns. comprising compds.

I [A = (un)saturated and/or (un)substituted 4-8 membered monocyclic or 7-12 membered bicyclic ring, containing 1 to 5 heteroatoms selected from the group consisting of O, N or S; ring A may further contain a carboxamide, sulfone, sulfonamide, or an acyl group; A1 = (un)saturated and/or (un) substituted 5-9 membered monocyclic or 8-14 membered polycyclic heterocycle containing at least one N; or A1 = substituted urea, iminourea or thiourea alicyclic or cyclic analog; Z1 = CH2, CH2O, O, NH, CO, S, SO, CHOH, SO2; Z2 = (un) substituted 1-5 carbon linker optionally containing one or more heteroatoms; alternatively Z1-Z2 may contain a carboxamide, sulfone, sulfonamide, alkenyl, acyl group, or aryl or heteroaryl ring; X = CO, SO2, S, O, substituted amine, substituted CH; Y = CO, SO2, substituted amine, etc.; Y5 = C or N; Y3 and Y4 independently = H, halo, (un) substitutedalkyl, -aryl, -alkene, etc.; or Y3 and Y4 together form a (un)saturated and/or (un) substituted 3-8 membered monocyclic or a 7-11 membered bicyclic ring optionally containing heteroatoms; or X and Y3 form a 3-7 membered monocyclic ring optionally containing heteroatoms; Rb = OH, alkoxy, arylamine, etc.], or a pharmaceutically acceptable salt thereof, methods of selectively inhibiting or antagonizing the avß3 and/or the  $\alpha \nu \beta 5$ integrin without significantly inhibiting the avß6 integrin, and methods to prepare I. Thus, e.g., II was prepared in four steps with oxadiazole ring forming via cyclization reaction of amide oxime III with cyclic anhydride IV (preparation given). I antagonize  $\alpha v \beta 3$ integrin with an IC50 values ranging from 0.1 nM to 100  $\mu M$  in the 293-cell assay. Similarly, I also antagonized integrin with an IC50 values of  $< 50 \mu M$  in the cell adhesion assay.

L67 ANSWER 4 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 5

ACCESSION NUMBER: 2001:923768 HCAPLUS

DOCUMENT NUMBER: 136:53681

TITLE: Preparation of cycloalkylalkanoic acids as

integrin receptor antagonists

INVENTOR (S): Khanna, Ish Kumar; Clare, Michael; Gasiecki,

Alan F.; Rogers, Thomas; Chen, Barbara;

Russell, Mark; Lu, Hwang-Fun Pharmacia Corporation, USA

PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 171 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIND DATE				APPL	ICAT		DATE							
						-													
WO 2001096307				A2 20011220				,	WO 2	001-	20010615								
WO	2001	0963	07		`A3		2002	0020815											
	W :	ΑE,	ΑG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,		
																GH,			
																LR,			
																PT,			
																US,			
								BY,							•	•	•		
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,		
																TR,			
								GN,									•		
ΕP			A2		2003	0312	1	EP 2	001-	9483	53		20010615						
	R:	AT,	BE,												SE,	MC,	PT,		
								MK,							•	•	•		

JP 2002-510450

US 2003-311299

20010615

20030821

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US 2000-211781P
                                                                P 20000615
                                            WO 2001-US19104
                                                                W 20010615
OTHER SOURCE(S):
                         MARPAT 136:53681
ED
     Entered STN: 21 Dec 2001
     The preparation of compds. [I; A = heteroaryl (e.g., pyridine, imidazole,
AB
     thiazole, oxazole, benzimidazole, imidazopyridine, etc.); n = 0-2, etc.;
     R1 = H, alkyl, etc.; R2, R3, R4, R5 = alkyl, alkoxy, etc.], their
     pharmaceutically acceptable salts and compns., and methods of selectively
     inhibiting or antagonizing the av83 and/or
             integrin, are described. Thus, a multi-step
     synthesis of the trifluoroacetate salt of 2-[4-[3-(2-
    pyridinylamino)propoxy]phenyl]cyclopropaneacetic acid (II) is given.
     Administration of I inhibits angiogenesis, tumor metastasis,
     tumor growth, osteoporosis, Paget's disease, humoral hypercalcemia of
    malignancy, retinopathy, macular degeneration, arthritis, periodontal
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20040819

20040304

L67 ANSWER 5 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN

atherosclerosis, and viral diseases.

T2

A1

ACCESSION NUMBER:

JP 2004525069

US 2004043988

PRIORITY APPLN. INFO.:

2004:566608 HCAPLUS

disease, smooth muscle cell migration, including restenosis and

DOCUMENT NUMBER:

INVENTOR(S):

141:123621

TITLE:

Preparation of pyrazole derivatives as

integrin receptor antagonists Penning, Thomas D.; Khilevich, Albert; Chen, Barbara B.; Gandhi,

Preete; Wang, Yaping; Downs,

Victoria; Boys, Mark L.; Russell, Mark;

Spangler, Dale P.; Huff, Renee M.

PATENT ASSIGNEE(S):

SOURCE:

Pharmacia Corporation, USA PCT Int. Appl., 113 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	PATENT NO.					KIND DATE			APPLICATION NO.						DATE			
WO	WO 2004058761			A1										- 2	0031	710		
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ.	BA.	BB.	BG	BR	BW	ВV	D7	C2	CII	
•		CN,	co,	CR,	CU.	CZ.	DE,	DK.	DM.	DZ,	EC,	EF.	EG,	EC,	ET,	CA,	CH,	
		GE,	GH.	GM.	HR.	HU.	·ID,	TI.	TN	IS,	.TD	KE,	KC,	ED,	ET,	GD,	GD,	
		LK.	LR.	LS.	LT.	T.II	LV,	MΔ,	MD,	MC,	ME,	MNI	MU	MY,	KK,	KZ,	LC,	
		NZ.	OM.	PG.	PH.	PI.	PT,	PO	, עניו	ec,	en,	eн,	EC,	PIA,	MZ,	NI,	NO,	
		TM.	TN.	TR	TT.	TZ.	UA,	IIG	no,	117	υC,	DE,	3G,	SK,	ъь,	SY,	TJ,	
	RW:	RW.	GH,	GM.	KE,	I.C	MW.	MZ	CD,	OZ,	VC,	VIV ,	YU,	ZA,	ZM,	ZW		
		BV	KC.	K7	MD	DII,	MW,	ma,	aυ,	эь,	52,	12,	UG,	ZM,	ZW,	AM,	AZ,	
		EC,	ET	ED,	, כנויו	RU,	TJ,	IM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	
		TO,	LI,	rk,	GB,	GR,	HU,	IE,	IT,	ьU,	MC,	ΝL,	PT,	RO,	SE,	SI,	SK,	
CA	25076	TE'	Dr,	вυ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG
	25079				AA		2004	0715	(	CA 20	003-2	25079	958		20	00312	219	
AU	20032	29740	)9		A1		2004	0722	1	AU 20	003-2	2974(	)9 ,		20	00312	219	
US	20050	0420	00		A1		2005(	0106	1	JS 20	003-1	74186	50		2.0	00311	719	
EP	15726	591			A1		2005	914	]	EP 20	003-8	31422	27		20	0312	219	
	R:	AT,	ΒE,	CH,	DΕ,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE.	MC.	PT.	
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ.	EE.	HU.	SK	·	
BR	20030	01687	75		Α	:	2005	1025	I	3R 20	003-1	16875	5		20	0312	119	
JP 2006513218				T2	:	20060	0420	BR 2003-16875 JP 2004-563834				20031219						
PRIORITY	PRIORITY APPLN. INFO.:			•										20				
											· - ·			•	. 20	12	. 20	

WO 2003-US40630 W 20031219 OTHER SOURCE(S): MARPAT 141:123621 Entered STN: 15 Jul 2004 Title compds. I [wherein M1 = heteroaryl, acyl, (un) substituted AB hydrocarbyl; R1 = O, CO, SOm, NHSO2, SO2NH, (un) substituted methylene or amino; m = 0-2; R4 = C or N; R5 = H, halo, (un) substituted hydrocarbyl, heteroaryl; R6 = an electron pair when R4 is nitrogen, or H, halo, (un) substituted hydrocarbyl, heterocyclo; or R4R5R6 = mono or bicyclic ring; X1 = 0, CH2, CH2O, NH, CO, SOm, CH(OH), alkenyl, alkynyl; X2 = (un)substituted linker; X3 = heterocyclic; Z1 = H, HO, cyano,
(un)substituted hydrocarbyl, heteroaryl; and pharmaceutically acceptable salts thereof) were prepared as integrin receptor antagonists. For example, 3-(1,3-benzodioxol-5-yl)-4-[1-methyl-5-[2-(5,6,7,8-tetrahydro-1,8-naphthyridin-2-yl)ethoxy]-1H-pyrazol-3-yl] butanoic acid (II) was given in a multiple-step synthesis starting from 2-aminonicotinaldehyde. The prepared title compds. I were tested for inhibition of  $\alpha v \beta 3$  and/or  $\alpha v \beta 5$ integrin receptors. Thus, I and their pharmaceutical compns. are useful for the treatment or prevention of conditions mediated by the  $\alpha v\beta 3$  or  $\alpha v\beta 5$  integrin receptor, such as tumor metastasis, solid tumor growth, angiogenesis, osteoporosis, humoral hypercalcemia of malignancy, smooth muscle cell migration, restenosis, atheroscelorosis, macular degeneration, retinopathy, and arthritis (no data). REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L67 ANSWER 6 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:566607 HCAPLUS DOCUMENT NUMBER: 141:123614 TITLE: Preparation of thiazoles, in particular thiazole butanoic acid derivatives, as integrin receptor antagonists INVENTOR(S): Wendt, John A.; Stenmark, Heather; Wu, Lisa; Wang, Yaping; Chen, Barbara B.; Penning, Thomas D.; Downs, Victoria; Boys, Mark L.; Russell, Mark; Spangler, Dale P. PATENT ASSIGNEE(S): Pharmacia Corporation, USA SOURCE: PCT Int. Appl., 110 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA	TENT	NO.			KIN	D :	DATE			APPL	ICAT	ION I	NO.		D	ATE			
WO	2004	0587	60		Al					WO 2003-US40629					20031219				
	W :	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,		
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,		
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,		
								RO,											
		TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW			
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG;	ZM,	ZW,	AM,	AZ,		
		BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,		
		ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,		
								CM,										TG	
CA 2510084								0715											

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AU 2003297408
                                 20040722
                           A1
                                              AU 2003-297408
                                                                     20031219 .
     US 2005004189
                           Α1
                                 20050106
                                              US 2003-741056
                                                                     20031219
     EP 1572690
                                 20050914
                                              EP 2003-814226
                           Α1
                                                                     20031219
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
     BR 2003016876
                           Α
                                 20051025
                                              BR 2003-16876
                                                                     20031219
PRIORITY APPLN. INFO.:
                                              US 2002-435030P
                                                                     20021220
                                              WO 2003-US40629
                                                                  W 20031219
OTHER SOURCE(S):
                          MARPAT 141:123614
     Entered STN: 15 Jul 2004
ED
     Title compds. I [wherein R1 = CHR2, NR3, O, S, SO2, NHSO2, SO2NH, C(:O);
AB
     R2 = H, OH, (un) substituted hydrocarbyl or alkoxy; R3 = H, (un) substituted
     hydrocarbyl, heteroaryl, or acyl; R4 = C, N; R5 = H, halo, (un)substituted
     hydrocarbyl, heteroaryl; R6 = electron pair when R4 = H; or R6 = H, halo,
     or (un) substituted hydrocarbyl; R7 = OH and derivs., SH and derivs., NH2
     and derivs., etc.; Z1 = H, OH, CN, (un) substituted hydrocarbyl,
     heteroaryl; and pharmaceutically acceptable salts thereof] were prepared for
     selectively inhibiting or antagonizing the
     \alpha v\beta 3 and/or \alpha v\beta 5
                           integrins (vitronectin
     receptors). For example, condensation of 4-(1-methyl-1,2,3,4-
     tetrahydropyrido[2,3-b]pyrazin-6-yl)butanethioamide (11-step synthesis
     given) with Et 3-(1,3-benzodioxol-5-yl)-6-chloro-5-oxohexanoate in dioxane
     at reflux, followed by saponification of the in-situ formed ester using NaOH in
     EtOH, gave II. Selected \alpha v \beta 3 and/or \alpha v \beta 5
     integrin antagonists I displayed AUC/oral dose ratios in
     the range of 1.8 - 6.8 when administered to rats. Thus, I and their
     pharmaceutical compns. are useful for the treatment of tumor metastasis,
     solid tumor growth, angiogenesis, osteoporosis, humoral hypercalcemia of
     malignancy, smooth muscle cell migration, restenosis, atherosclerosis,
     macular degeneration, retinopathy, and arthritis (no data).
L67 ANSWER 7 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
                          2004:859471 HCAPLUS
DOCUMENT NUMBER:
                          143:115774
TITLE:
                          Synthesis and tissue distribution of 14C-nonpeptide
                         αvβ3 antagonists
AUTHOR (S):
                         McKinnis, Bradley R.; Albin, Lesley A.; Doom, James
                          P.; Gasiecki, Alan F.; Hotz, Kathy J.; Keene, Jeffery
                          L.; Khanna, Ish K.; Kraus, Lori J.; Liley,
                         Matt R.; Likos, John J.; Nagarajan, Srinivasan
                          ; Rogers, Thomas E.; Singh, Rajendra K.
CORPORATE SOURCE:
                         Pfizer Global Research and Development, St. Louis, MO,
SOURCE:
                         Synthesis and Applications of Isotopically Labelled
                         Compounds, Proceedings of the International Symposium,
                         8th, Boston, MA, United States, June 1-5, 2003 (2004),
                         Meeting Date 2003, 413-416. Editor(s): Dean, Dennis
                         C.; Filer, Crist N.; McCarthy, Keith E. John Wiley &
                         Sons Ltd.: Chichester, UK.
                         CODEN: 69FZAZ; ISBN: 0-470-86365-X
DOCUMENT TYPE:
                         Conference
LANGUAGE:
                         English
OTHER SOURCE(S):
                         CASREACT 143:115774
     Entered STN: 18 Oct 2004
AB
     Three peptidomimetic integrin av 33
     antagonists, I (R = Me, 3-pyridyl; X = 14CH2) and II (X = 14CH2),
     are prepared; their antagonism of integrin
     \alpha v \beta 3 and their distributions in the tissues of rats are determined
     14C-labeled 2-(3-hydroxypropylamino)pyridine-1-oxide is prepared from
     2-chloropyridine N-oxide hydrochloride and 3-amino-3-[14C]-aminopropanol;
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Mitsunobu coupling with p-hydroxyphenyl-substituted esters, redn . of the N-oxide, and ester hydrolysis yields I (R = Me, 3-pyridyl; X = 14CH2) and II (X = 14CH2). I (R = Me; X = 14CH2) is generated in an attempted preparation of II (X = 14CH2); reduction of the N-oxide ester intermediate in the preparation of II (X = 14CH2) with palladium on carbon and cyclohexene in refluxing isopropanol leads to reduction of the N-oxide and cleavage of the cyclopropane ring rather than reduction of the N-oxide alone. Reduction of the N-oxide ester intermediate in the preparation of II (X = 14CH2) with triphenylphosphine and iron in refluxing acetic acid reduces the N-oxide without cleaving the cyclopropane. I (R = Me, 3-pyridyl; X = 14CH2) and II (X = 14CH2) are obtained with radiochem. purities of  $\geq$  97.4%. Tissue distribution, mass balance, and clearance studies for I (R = Me, 3-pyridyl; X = 14CH2) and II (X = 14CH2) are performed.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L67 ANSWER 8 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:923795 HCAPLUS

DOCUMENT NUMBER: 136:53749

TITLE: Preparation of heteroarylalkanoic acids as

integrin receptor antagonists

INVENTOR(S): Nagarajan, Scrinivasan Raj; Khanna, Ish

Kumar; Tollefson, Michael B.;
Mohler, Scott B.; Chen, Barbara;
Russell, Mark; Devadas, Balekudru;
Penning, Thomas D.; Schretzman, Lori
A.; Spangler, Dale P.; Boys, Mark
Laurence; Chandrakumar, Nizal Samuel;

Lu, Hwang-Fun

PATENT ASSIGNEE(S): Pharmacia Corporation, USA

SOURCE: PCT Int. Appl., 368 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENI	NO.	KIN	JD DATE	;	A	PPLIC	CATION	NO.		D	ATE	
	1096334 1096334			1220	W	0 200	01-US19	375		2	0010	615
<b>W</b> :	AE, AG, CO, CR, HR, HU, LT, LU, RU, SD, VN, YU, Y: GH, GM, DE, DK,	AL, AM, CU, CZ, ID, IL, LV, MA, SE, SG, ZA, ZW, KE, LS, ES, FI,	AT, AU, DE, DK, IN, IS, MD, MG, SI, SK, AM, AZ,	AZ, DM, JP, MK, SL, BY, SD, GR,	DZ, KE, MN, TJ, KG, SL, IE,	EE, H KG, H MW, N TM, T KZ, N SZ, T IT, I	ES, FI, KP, KR, MX, MZ, FR, TT, MD, RU, FZ, UG, LU, MC,	GB, KZ, NO, TZ, TJ, ZW, NL,	GD, LC, NZ, UA, TM AT, PT,	GE, LK, PL, UG, BE, SE,	GH, LR, PT, US,	GM, LS, RO, UZ,
	2133023				. U	S 200	01-8819	13		20	0010	515
EP 128 R: JP 200	9983 AT, BE, IE, SI, 4511434 4092497	CH, DE, LT, LV, T2 A1	2 2003 DK, ES, FI, RO, 2 2004	FR, MK, 0415	GB, G CY, J U	GR, ] AL, T P 200 S 200 S 200		LU, 76 85 81P	NL,	SE, 20 20 20	00106 MC, 00106 00309	PT, 515 905 515

WO 2001-US19375 W 20010615

OTHER SOURCE(S): MARPAT 136:53749

Entered STN: 21 Dec 2001 ED

AB Title compds. A1Z2Z1AXYY5(Y3)(Y4)CH2CORb [I; wherein ring A = (un) substituted 4-8 membered monocyclic or 7-12 membered bicyclic ring containing 1-4 heteroatoms, selected from O, N, or S; A1 = (un)substituted 5-9 membered monocyclic or 7-14 membered polycyclic heterocycle containing at least 1 N and optionally 1-4 heteroatoms or groups selected from O, N, S, SO2, or CO; Z1 = CH2, O, CH2O, NH, CO, S, SO, CH(OH), and SO2; Z2 = (un) substituted 1-5 C linker optionally containing 1 or more heteroatoms selected from O, S, and N; Z1Z2 may contain a carboxamide, sulfone, sulfonamide, alkenyl, alkynyl, acyl, or (un)substituted 5- or 6-membered (hetero) aryl; X = CHRe, NRf, O, S, SO2, or CO; Re = H, (cyclo) alkyl, alkoxy(alkyl), OH, alkynyl, alkenyl, haloalkyl, thioalkyl, or aryl; Rf = H, (halo)alkyl, aryl, or benzyl; Y = (CH2)p, CHRg, NRg, CO, or SO2; Rg = H, (halo)alkyl, alkoxyalkyl, alkynyl, (hetero)aryl, OH, alkoxy, or carboxyalkyl; p = 0-1; XY may contain acyl, alkyl, sulfonyl, amino, (thio)ether, carboxamido, sulfonamido, aminosulfonyl, or olefin; Y3 and Y4 = independently H, (halo)alkyl, halo, (hetero)aryl, hydroxyalkyl, alkynyl, etc.; Rb = X2Rh; X2 = O, S, or NRj; Rh and Rj = independently H, (ar) alkyl, acyl, or alkoxyalkyl; with provisos] and their pharmaceutically acceptable salts were prepared for selectively antagonizing the  $\alpha v\beta 3$  and/or the  $\alpha v\beta 5$ integrin without significantly antagonizing the fibrinogen IIb/IIIa integrin. For example, 3-(hydroxymethyl)benzonitrile was protected with 3,4-dihydro-2H-pyran (89%) and treated with HONH2-HCl to give the benzenecarboximidamide (98%). Cyclization with 3-methylglutaric anhydride in the presence of MeI (64%) and deprotection (98%) gave the Me 1,2,4-oxadiazolebutanoate (64%). Oxidation to the aldehyde, followed by reductive addition of 2-aminopyridine and workup, afforded the oxadiazolebutanoic acid (II). In vitronectin adhesion assays, I antagonized the ανβ3 integrin and the  $\alpha v \beta 5$ integrin with IC50 values of 0.1 nM to 100  $\mu M$  and < 50  $\mu M$ , resp. I are useful for the treatment of tumor metastasis, solid tumor growth, angiogenesis, osteoporosis, humoral hypercalcemia of malignancy, smooth muscle cell migration, restenosis, atherosclerosis, macular degeneration, retinopathy, and arthritis (no data).

L67 ANSWER 9 OF 11 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:923771 HCAPLUS

DOCUMENT NUMBER: 136:53683

TITLE:

Preparation of dihydrostilbene alkanoic acid derivatives useful as vitronectin antagonists INVENTOR (S): Rogers, Thomas; Clare, Michael; Fun Lu, Hwang;

Russell, Mark; Malecha, James W.; Khanna, Ish

Kumar; Penning, Thomas; Nagarajan,

Srinivasan Raj

PATENT ASSIGNEE(S): Pharmacia Corporation, USA

PCT Int. Appl., 163 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001096310	A1	20011220	WO 2001-US19330	20010615
W: AE, AG, AL,	AM, AT	, AU, AZ, BA	, BB, BG, BR, BY, BZ,	CA, CH, CN,

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CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
              HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
              LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
              RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
              VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
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     US 2001052500
                                              US 2001-882647
                           A1
                                 20011220
                                                                      20010615
                           A5
     AU 2001068490
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     US 2002099209
                                              US 2001-882137
                           A1
                                 20020725
                                                                      20010615
     US 6720315
                           B2
                                 20040413
     EP 1289959
                           A1
                                 20030312
                                              EP 2001-946439
                                                                      20010615
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     JP 2004503540
                           T2
                                 20040205
                                              JP 2002-510453
                                                                      20010615
     US 6833366
                           B1
                                 20041221
                                              US 2003-657932
                                                                      20030909
PRIORITY APPLN. INFO.:
                                              US 2000-211780P
                                                                   P 20000615
                                              US 2001-882137
                                                                   A3 20010615
                                              WO 2001-US19330
                                                                   W 20010615
OTHER SOURCE(S):
                          MARPAT 136:53683
     Entered STN:
                    21 Dec 2001
AB
     The preparation of [I; wherein the "A ring" = 4-8 membered monocyclic, or 7-12
     membered bicyclic heteroarene; A1 = 5-9 membered monocyclic, or 7-12
     membered polycyclic heterocycle; Z1 = CH2, CH2O, O, NH, CO, S, etc.; Z2 =
     1-5 carbon linker optionally substituted with O, S, or N; X = alkyl, O,
     amino, CO, etc.; Y = substituted C; Ra = H, alkyl, alkenyl, etc.; R1 = H,
     alkyl, hydroxy, etc.; R2 = H, alkyl, etc.; R3 = H, alkyl, halogen, etc.],
     or a pharmaceutically acceptable salt or composition thereof, and methods of
     selectivelyavß3
                      inhibiting or antagonizing
     the \alpha \nu \beta 3 and/or the \alpha \nu \beta 5
                                     integrin,
     are described. Thus, a multi-step preparation of 3-[[3-(2-
     pyridinylamino)propoxy]phenyl]propanoic acid II was given. Administration
     of I inhibits angiogenesis, tumor metastasis, tumor growth,
     osteoporosis, Paget's disease, humoral hypercalcemia of malignancy,
     retinopathy, macular degeneration, arthritis, periodontal disease, smooth
     muscle cell migration, including restenosis and atherosclerosis, and viral
     diseases.
REFERENCE COUNT:
                          3
                                THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 10 OF 11 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on
     STN
                                                          DUPLICATE 4
ACCESSION NUMBER:
                    2006:29662 BIOSIS
DOCUMENT NUMBER:
                    PREV200600040384
TITLE:
                    Dihydrostilbene alkanoic acid derivatives.
AUTHOR(S):
                    Rogers, Thomas [Inventor]; Clare, Michael [Inventor];
                    Lu, Hwang-Fun [Inventor]; Russell, Mark [Inventor];
                    Malecha, James W. [Inventor]; Khanna, Ish Kumar
                     [Inventor]; Penning, Thomas [Inventor];
                    Nagarajan, Srinivasan Raj [Inventor]
CORPORATE SOURCE:
                    Ballwin, MO USA
                    ASSIGNEE: Pharmacia Corporation
PATENT INFORMATION: US 06833366 20041221
SOURCE:
                    Official Gazette of the United States Patent and Trademark
                    Office Patents, (DEC 21 2004)
                    CODEN: OGUPE7. ISSN: 0098-1133.
DOCUMENT TYPE:
                    Patent
LANGUAGE:
                    English
ENTRY DATE:
                    Entered STN: 28 Dec 2005
```

Last Updated on STN: 28 Dec 2005

Entered STN: 28 Dec 2005 ED

Last Updated on STN: 28 Dec 2005

The present invention relates to a class of compounds represented by the AB Formula 1. or a pharmaceutically acceptable salt thereof, pharmaceutical compositions comprising compounds of the Formula 1, and methods of selectively inhibiting or antagonizing the alpha(v)beta(3) and/or the alpha(v)beta(5) integrin.

ANSWER 11 OF 11 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

ACCESSION NUMBER:

2004:257549 BIOSIS

DOCUMENT NUMBER:

PREV200400257507

Dihydrostilbene alkanoic acid derivatives.

AUTHOR (S):

Rogers, Thomas [Inventor, Reprint Author]; Clare, Michael

[Inventor]; Lu, Hwang-Fun [Inventor]; Russell,

Mark [Inventor]; Malecha, James W. [Inventor]; Khanna,

Ish Kumar [Inventor]; Penning, Thomas

[Inventor]; Nagarajan, Srinivasan Raj [Inventor];

Stenmark, Heather [Inventor]

CORPORATE SOURCE:

Manchester, MO, USA

ASSIGNEE: Pharmacia Corporation

PATENT INFORMATION: US 6720315 20040413

SOURCE:

Official Gazette of the United States Patent and Trademark

Office Patents, (Apr 13 2004) Vol. 1281, No. 2. http://www.uspto.gov/web/menu/patdata.html. e-file.

ISSN: 0098-1133 (ISSN print).

DOCUMENT TYPE:

Patent English

LANGUAGE: ENTRY DATE:

Entered STN: 12 May 2004

Last Updated on STN: 12 May 2004

Entered STN: 12 May 2004

Last Updated on STN: 12 May 2004

The present invention relates to a class of compounds represented by the AΒ Formula 1. ##STR1## or a pharmaceutically acceptable salt thereof, pharmaceutical compositions comprising compounds of the Formula 1, and methods of selectively inhibiting or antagonizing the alphaVbeta3 and/or the alphaVbeta5 integrin.

=> file stnquide

FILE 'STNGUIDE' ENTERED AT 13:02:32 ON 30 MAY 2006 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY, JAPAN SCIENCE AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: May 26, 2006 (20060526/UP).

L8

L9

L10

(FILE 'HOME' ENTERED AT 11:52:34 ON 30 MAY 2006)

FILE 'STNGUIDE' ENTERED AT 11:52:41 ON 30 MAY 2006

FILE 'ZCAPLUS' ENTERED AT 11:57:50 ON 30 MAY 2006 E US2003-743354/APPS

FILE 'HCAPLUS' ENTERED AT 11:58:09 ON 30 MAY 2006
L1 1 SEA ABB=ON PLU=ON US2003-743354/APPS
SAVE TEMP L1 GEM354HCAAPP/A

FILE 'STNGUIDE' ENTERED AT 11:58:26 ON 30 MAY 2006

FILE 'HCAPLUS' ENTERED AT 11:58:33 ON 30 MAY 2006 D SCAN

FILE 'STNGUIDE' ENTERED AT 11:58:39 ON 30 MAY 2006

FILE 'STNGUIDE' ENTERED AT 11:59:20 ON 30 MAY 2006

FILE 'REGISTRY' ENTERED AT 12:00:49 ON 30 MAY 2006

FILE 'HCAPLUS' ENTERED AT 12:00:52 ON 30 MAY 2006
L3 TRA PLU=ON L1 1- RN : 387 TERMS

FILE 'REGISTRY' ENTERED AT 12:00:55 ON 30 MAY 2006 L4 387 SEA ABB=ON PLU=ON L3 SAVE TEMP L4 GEM354REGAPP/A

FILE 'STNGUIDE' ENTERED AT 12:01:16 ON 30 MAY 2006

FILE 'REGISTRY' ENTERED AT 12:01:30 ON 30 MAY 2006
L5 131 SEA ABB=ON PLU=ON L4 AND ?NAPHTHYRIDIN?/CNS
L6 80 SEA ABB=ON PLU=ON L5 AND ?OXADIAZOL?/CNS
L7 1 SEA ABB=ON PLU=ON L6 AND ?IODO?/CNS
D SCAN

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FILE 'REGISTRY' ENTERED AT 12:06:06 ON 30 MAY 2006

1 SEA ABB=ON PLU=ON L6 AND I/ELS
1 SEA ABB=ON PLU=ON L5 AND I/ELS
6 SEA ABB=ON PLU=ON L4 AND I/ELS
D SCAN

SAVE TEMP L9 GEM354ES/A D SCAN L9

FILE 'STNGUIDE' ENTERED AT 12:08:14 ON 30 MAY 2006 D QUE STAT L9

FILE 'REGISTRY' ENTERED AT 12:09:19 ON 30 MAY 2006 D IDE L9

FILE 'STNGUIDE' ENTERED AT 12:09:20 ON 30 MAY 2006

FILE 'HCAPLUS, TOXCENTER, USPATFULL' ENTERED AT 12:11:11 ON 30 MAY 2006
L11 3 SEA ABB=ON PLU=ON L9
D SCAN
SAVE TEMP L11 GEM354MULS1/A

FILE 'STNGUIDE' ENTERED AT 12:11:53 ON 30 MAY 2006 D SAVED

FILE 'LREGISTRY' ENTERED AT 12:12:27 ON 30 MAY 2006 L12 STR 724770-27-8

FILE 'BEILSTEIN' ENTERED AT 12:13:09 ON 30 MAY 2006 L13 0 SEA SSS FUL L12 SAVE TEMP L13 GEM354BEI1/A D QUE STAT

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L17 0 SEA ABB=ON PLU=ON L16 NOT L9

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FILE 'CHEMINFORMRX' ENTERED AT 12:18:59 ON 30 MAY 2006
L19 0 SEA SSS SAM L14 ( 0 REACTIONS)
D QUE STAT

L20 0 SEA SSS FUL L14 ( 0 REACTIONS) SAVE TEMP L20 GEM354CHM1/A

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FILE 'MARPAT' ENTERED AT 12:20:15 ON 30 MAY 2006 L21 0 SEA SSS SAM L14 D QUE STAT

L22 0 SEA SSS FUL L14 SAVE TEMP L22 GEM354MAR1/A

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FILE 'STNGUIDE' ENTERED AT 12:23:45 ON 30 MAY 2006

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FILE 'WPIX' ENTERED AT 12:24:13 ON 30 MAY 2006
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L25
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L26
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L27
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L28
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                D QUE STAT
L29
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L30
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L32
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                QUE ABB=ON PLU=ON SCHRETZMAN, L?/AU
L34
                QUE ABB=ON PLU=ON TOLLEFSON, M?/AU
L35
                QUE ABB=ON PLU=ON CHANDRAKUMAR, N?/AU
L36
                QUE ABB=ON PLU=ON CHANDRAKOMAK,
QUE ABB=ON PLU=ON KHANNA, I?/AU
QUE ABB=ON PLU=ON NGUYEN, M?/AU
QUE ABB=ON PLU=ON DOWNS, V?/AU
QUE ABB=ON PLU=ON MOHLER, S?/AU
L37
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L38 L39 L40

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L41
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L42
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L43
L44
               QUE ABB=ON PLU=ON WANG, Y?/AU
L*** DEL
               QUE KHILEVICH, A?/U
L45
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L46
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L47
               OUE ABB=ON PLU=ON YU, Y?/AU
L48
               QUE ABB=ON PLU=ON WENDT, J?/AU
               QUE ABB=ON PLU=ON STENMARK, H?/AU
L49
L50
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L51
               QUE ABB=ON PLU=ON HUFF, R?/AU
L52
               QUE ABB=ON PLU=ON NAGARAJAN, S?/AU
L53
               QUE ABB=ON PLU=ON DEVADAS, B?/AU
L54
               QUE ABB=ON PLU=ON LU, H?/AU
               OUE ABB=ON PLU=ON RUSSEL, M?/AU
L55
               OUE ABB=ON PLU=ON SPANGLER, D?/AU
L56
               OUE ABB=ON PLU=ON PARIKH, M?/AU
L57
               OUE ABB=ON PLU=ON PHARMACIA/PA,CS,SO
L58
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FILE 'STNGUIDE' ENTERED AT 12:34:29 ON 30 MAY 2006

FILE 'HCAPLUS, MEDLINE, BIOSIS, PASCAL, JICST-EPLUS, CABA, LIFESCI, DRUGU, DRUGB, VETU, VETB, WPIX, SCISEARCH, CONF, CONFSCI, DISSABS' ENTERED AT 12:37:36 ON 30 MAY 2006

- L59 194315 SEA ABB=ON PLU=ON (L33 OR L34 OR L35 OR L36 OR L37 OR L38 OR L39 OR L40 OR L41 OR L42 OR L43 OR L44 OR L45 OR L46)
- L60 83410 SEA ABB=ON PLU=ON (L47 OR L48 OR L49 OR L50 OR L51 OR L52 OR L53 OR L54 OR L55 OR L56 OR L57)
- L61 904 SEA ABB=ON PLU=ON (L59 OR L60) AND ?INTEGRIN?
- L62 10520 SEA ABB=ON PLU=ON (L33 OR L34 OR L35 OR L36 OR L37) OR (L39 OR L40 OR L41 OR L42) OR (L45 OR L46) OR (L48 OR L49) OR (L51 OR L52 OR L53) OR (L55 OR L56 OR L57)
- L63 215 SEA ABB=ON PLU=ON (L38 OR L43 OR L44 OR L47 OR L50 OR L54)
  AND L58
- L64 94 SEA ABB=ON PLU=ON (L62 OR L63) AND ?INTEGRIN?
- L65

  85 SEA ABB=ON PLU=ON L64 AND (?INTEGRIN?(L)(?ANTAGON? OR
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  INTERRUPT? OR CONTROL? OR MODERAT? OR MODULAT? OR ?REGULAT? OR
  ?PREVENT? OR ?REDUC? OR ?IMPED? OR ?SUPPRESS? OR REPRESS? OR
  RETARD? OR SLOW?))
- L66 21 SEA ABB=ON PLU=ON L65 AND (ALKANOIC? OR HETEROALKANOIC? OR ?BUTANOIC?)
- L67

  11 DUP REM L66 (10 DUPLICATES REMOVED)

  ANSWERS '1-9' FROM FILE HCAPLUS

  ANSWERS '10-11' FROM FILE BIOSIS

  SAVE TEMP L67 GEM354MULINV/A

FILE 'STNGUIDE' ENTERED AT 12:57:16 ON 30 MAY 2006

- D SAVED
- D QUE STAT L16
- D QUE NOS L17
- D QUE NOS L24
- D QUE STAT L18
- D QUE STAT L20
- D QUE STAT L22
- D QUE STAT L32
- D QUE STAT L11

FILE 'HCAPLUS, TOXCENTER, USPATFULL, WPIX' ENTERED AT 13:00:18 ON 30 MAY

2006

L68

2 DUP REM L11 L18 L20 L22 L32 (2 DUPLICATES REMOVED)
ANSWER '1' FROM FILE HCAPLUS
ANSWER '2' FROM FILE USPATFULL

FILE 'STNGUIDE' ENTERED AT 13:00:24 ON 30 MAY 2006

FILE 'HCAPLUS, USPATFULL' ENTERED AT 13:00:34 ON 30 MAY 2006 D IBIB ED AB IND HITSTR

FILE 'STNGUIDE' ENTERED AT 13:00:35 ON 30 MAY 2006

FILE 'HCAPLUS, USPATFULL' ENTERED AT 13:00:45 ON 30 MAY 2006 D IBIB AB HITSTR 2

FILE 'STNGUIDE' ENTERED AT 13:00:46 ON 30 MAY 2006 D QUE STAT L67

FILE 'HCAPLUS, BIOSIS' ENTERED AT 13:02:10 ON 30 MAY 2006 D IBIB ED AB L67 1-11

FILE 'STNGUIDE' ENTERED AT 13:02:12 ON 30 MAY 2006

FILE 'STNGUIDE' ENTERED AT 13:02:32 ON 30 MAY 2006

FILE HOME

FILE STNGUIDE FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: May 26, 2006 (20060526/UP).

#### FILE ZCAPLUS

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FILE COVERS 1907 - 30 May 2006 VOL 144 ISS 23 FILE LAST UPDATED: 29 May 2006 (20060529/ED)

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FILE COVERS 1907 - 30 May 2006 VOL 144 ISS 23 FILE LAST UPDATED: 29 May 2006 (20060529/ED)

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#### FILE WPIX

FILE LAST UPDATED: 26 MAY 2006 <20060526/UP>
MOST RECENT DERWENT UPDATE: 200634 <200634/DW>
DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

>>> FOR A COPY OF THE DERWENT WORLD PATENTS INDEX STN USER GUIDE, PLEASE VISIT:

http://www.stn-international.de/training\_center/patents/stn\_guide.pdf <

>>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE http://scientific.thomson.com/support/patents/coverage/latestupdates/

>>> PLEASE BE AWARE OF THE NEW IPC REFORM IN 2006, SEE http://www.stn-international.de/stndatabases/details/ipc\_reform.html and http://scientific.thomson.com/media/scpdf/ipcrdwpi.pdf <<<

#### FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 29 MAY 2006 HIGHEST RN 885947-35-3 DICTIONARY FILE UPDATES: 29 MAY 2006 HIGHEST RN 885947-35-3

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 6, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

FILE TOXCENTER

FILE COVERS 1907 TO 30 May 2006 (20060530/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

The MEDLINE file segment has been updated with 2006 MEDLINE data and features. See HELP RLOAD for details.

TOXCENTER thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

See http://www.nlm.nih.gov/mesh/

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\_med\_data\_changes.html http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\_2006\_MeSH.html for a description of changes.

FILE USPATFULL

FILE COVERS 1971 TO PATENT PUBLICATION DATE: 30 May 2006 (20060530/PD)
FILE LAST UPDATED: 30 May 2006 (20060530/ED)
HIGHEST GRANTED PATENT NUMBER: US7055175
HIGHEST APPLICATION PUBLICATION NUMBER: US2006112473
CA INDEXING IS CURRENT THROUGH 30 May 2006 (20060530/UPCA)
ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 30 May 2006 (20060530/PD)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Feb 2006
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Feb 2006

FILE LREGISTRY
LREGISTRY IS A STATIC LEARNING FILE

NEW CAS INFORMATION USE POLICIES, ENTER HELP USAGETERMS FOR DETAILS.

FILE BEILSTEIN
FILE LAST UPDATED ON MARCH 15, 2006

FILE COVERS 1771 TO 2006.
FILE CONTAINS 9,516,393 SUBSTANCES

>>>PLEASE NOTE: Reaction Data and substance data are stored in separate documents and can not be searched together in one query. Reaction data for BEILSTEIN compounds may be displayed immediately with the display codes PRE (preparations) and REA (reactions). A substance answer set retrieved after the search for a chemical name, a compounds with available reaction information by combining with PRE/FA, REA/FA or more generally with RX/FA. The BEILSTEIN Registry Number (BRN) is the link between a BEILSTEIN compound and belonging reactions. For mo detailed reaction searches BRNs can be searched as reaction partner BRNs Reactant BRN (RX.RBRN) or Product BRN (RX.PBRN).<<<

>>> FOR SEARCHING PREPARATIONS SEE HELP PRE <<<

\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

- \* PLEASE NOTE THAT THERE ARE NO FORMATS FREE OF COST.
- \* SET NOTICE FEATURE: THE COST ESTIMATES CALCULATED FOR SET NOTICE
- \* ARE BASED ON THE HIGHEST PRICE CATEGORY. THEREFORE; THESE
- \* ESTIMATES MAY NOT REFLECT THE ACTUAL COSTS.
- \* FOR PRICE INFORMATION SEE HELP COST

\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

NEW

- \* PATENT NUMBERS (PN) AND BABS ACCESSION NUMBERS (BABSAN) CAN NOW BE SEARCHED, SELECTED AND TRANSFERRED.
- \* NEW DISPLAY FORMATS ALLREF, ALLP AND BABSAN SHOW ALL REFERENCES, ALL PATENT REFERENCES, OR ALL BABS ACCESSION NUMBERS FOR A COMPOUND AT A GLANCE.

FILE CHEMINFORMRX

FILE LAST UPDATED: 8 MAR 2006 <20060308/UP>

FILE MARPAT

FILE CONTENT: 1961-PRESENT VOL 144 ISS 22 (20060526/ED)

SOME MARPAT RECORDS ARE DERIVED FROM INPI DATA FOR 1961-1987

MOST RECENT CITATIONS FOR PATENTS FROM MAJOR ISSUING AGENCIES (COVERAGE TO THESE DATES IS NOT COMPLETE):

```
2006062725 23 MAR 2006
DE 102004045029 16 MAR 2006
EP
        1634887 15 MAR 2006
JP
     2006073583 16 MAR 2006
WO
     2006045852 04 MAY 2006
GB
        2416167 18 JAN 2006
FR
        2875804 31 MAR 2006
RU
        2270725 27 FEB 2006
CA
        2518664 10 MAR 2006
```

Expanded G-group definition display now available.

New CAS Information Use Policies, enter HELP USAGETERMS for details.

#### FILE MEDLINE

FILE LAST UPDATED: 27 MAY 2006 (20060527/UP). FILE COVERS 1950 TO DATE.

On December 11, 2005, the 2006 MeSH terms were loaded.

The MEDLINE reload for 2006 is now (26 Feb.) available. For details on the 2006 reload, enter HELP RLOAD at an arrow prompt (=>). See also:

```
http://www.nlm.nih.gov/mesh/http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html http://www.nlm.nih.gov/pubs/techbull/nd05/nd05_med_data_changes.html http://www.nlm.nih.gov/pubs/techbull/nd05/nd05_2006_MeSH.html
```

OLDMEDLINE is covered back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BIOSIS

FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 24 May 2006 (20060524/ED)

FILE PASCAL

FILE LAST UPDATED: 29 MAY 2006 <20060529/UP>

FILE COVERS 1977 TO DATE.

>>> SIMULTANEOUS LEFT AND RIGHT TRUNCATION IS AVAILABLE IN THE BASIC INDEX (/BI) FIELD <<<

FILE JICST-EPLUS

FILE COVERS 1985 TO 30 MAY 2006 (20060530/ED)

THE JICST-EPLUS FILE HAS BEEN RELOADED TO REFLECT THE 1999 CONTROLLED TERM (/CT) THESAURUS RELOAD.

FILE CABA

FILE COVERS 1973 TO 3 May 2006 (20060503/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

The CABA file was reloaded 7 December 2003. Enter HELP RLOAD for details.

FILE LIFESCI

FILE COVERS 1978 TO 12 May 2006 (20060512/ED)

FILE DRUGU

FILE LAST UPDATED: 29 MAY 2006 <20060529/UP>

>>> DERWENT DRUG FILE (SUBSCRIBER) <<<

>>> FILE COVERS 1983 TO DATE <<<

>>> THESAURUS AVAILABLE IN /CT <<<

FILE DRUGB

>>> FILE COVERS 1964 TO 1982 - CLOSED FILE <<<

FILE VETU

FILE LAST UPDATED: 02 JAN 2002 <20020102/UP>

FILE COVERS 1983-2001

FILE VETB

FILE LAST UPDATED: 25 SEP 94 <940925/UP>

FILE COVERS 1968-1982

FILE SCISEARCH

FILE COVERS 1974 TO 25 May 2006 (20060525/ED)

SCISEARCH has been reloaded, see HELP RLOAD for details.

FILE CONF

FILE LAST UPDATED: 23 DEC 2005 <20051223/UP>

FILE COVERS 1976 TO 2005.

<>< CONF IS NO LONGER BEING UPDATED AS OF JANUARY 2006 >>>

FILE COVERS 1973 TO 10 Apr 2006 (20060410/ED)

CSA has resumed updates, see NEWS FILE

=>

FILE DISSABS FILE COVERS 1861 TO 25 MAY 2006 (20060525/ED)

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searched by D. Arnold 571-272-2532